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COMPLETE SPECIFICATION

The inventors of this invention in the sense of being the actual devisers thereof within the caning of Sex 16 of the Patents Act 1949 are:—Helmut Nonnenmacher, Hansmeaning of Sect 16 of the Patents Act 1949 are:—HELBOUT NUNESHOWARDERS, LEASE-MARTIN WEITZ and CALL-HEINZ WILLENDO, clitzens of the Federal Republic of Germany, residing, respectively at 9 Wolfmanstrasse, Ludwighhafen/Rhein; 20 Carl-Bosch-Ring, Frankenthal/Pfak; and 84 Wittelbachstrasse, Ludwigshafen/Rhein; Federal Republic of Germany.

Separation of Methacrylic Acid and Acrylic Acid

methocrytic acid and acrytic acid".

In numerous processes for the production of (meth)acrylic soid these compounds are obtained in impure form, especially as aqueous ornamen in impure form, especially as equieous 20 solutions which contain small amounts of saturand carboxylic acids. Purification is usually carried out by distribution. Since however (meth)acrylic acid has a higher boiling point than water, considerable amounts of heat

point than water, considerable amounts or next are required for separation if the solution has a high water content, owing to the high heat of vaporization of water. It is therefore advan-tageous to extract (meth)acrylic acid from aqueous solutions and to isolate it by distillation 30 of the organic phase.

A large number of solvents have been

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proposed for the extraction of (meth)acrylic acid from aqueous solutions: percoleum ether, benzene, tohuene, chloroform, carbon tetra-35 chloride, dichloroethame, trichloroethylene,

We, RADECHE AVILIN- & SOBA-FABUX
ANTIDUSERHLICHAY, is German Joint Stock
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is to be performed, to be particularly described
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service of the following structure of the following proposed passe (sulting-out-effect),
while many selvents necessitate the addition of
the recovery of the salt from the aqueeu
selvent proposed have low
the man "fat least one of the compounds
methocytic seid and scrylic scid and scrylic acid and scrylic scid and scrylic scid and scrylic acid and scrylic scid and scrylic scid. acid. 10 separate the solvent from the extract it must be distilled off; this means larger hear consumption. Some solvents are difficult to 45 separate from (meth) acrylic acid by distillation, while many solvents necessitate the addition of

Moreover, the solvents proposed have low partition coefficients $K(K=\frac{C_1}{C})$ where Cdenotes the concentration of (meth)scrylic acid in percent by weight in the organic phase and 55

C₂ denotes the concentration or (memory) acid in percent by weight in the aqueous phase). Low parthion coefficients necessing multi-coefficients necessing multidenotes the concentration of (meth)acrylic stage extraction or the use of larger amounts of solvent in order to ensure practically quantita-tive separation of (meth)acrylic acid from the

tire separation of (meth)acrylic acid from the aqueous phase. Furthermare, imputities other than water, perticularly prepiants acid and accid exid, cannot be removed astiriaconsily. It is an object of the invention to provide a process for separating (meth)acrylic exid, in which the solvent has a higher partition co-efficient than solvents which have inhirated been emclent than solvents which nave nather to been proposed, in which the (methlecytic acid is separated practically quantitatively in a few 70 certaction stages, in which the solvent need not be distilled, in which the boiling point of the solvent is higher than the boiling point of (methlecytic acid, in which the addition of a

salt to the aqueous phase is not necessary and in which the main impurities, i.e. proplants acid and acetic acid, can be easily separated.

These and other objects are achieved in a

answe and other objects are achieved in a 5 process for the separation of (meth)acrylic acid by extraction with a lacram having 4 to 7 ring members and beauting a hydrocarbon radical as substituent on the nitrogen atom.

The new processes countries of the control of the contr

The new process is suitable for separating (meth)acrylic acid from both liquid and vaporous mixtures containing (meth)acrylic acid. Aqueous solutions obtained in commercial processes and containing about 2 to 50%

15 preferably 5 to 30%, by weight of (meth)acryl ocid are particularly suitable. The aqueon solutions may also contain other water-so organic or inorganic substances, such as lower alcohols, saturated carboxytic acids, carbon 20 dioxide or metal sales. Experience has shown that such substances do not affect the extraction

that successiones so not ancer the extraction of (methacrylic acid even if they are also extracted by the solvent. It is particularly advantageous to use aqueous (methacrylic acid solutions obtained in the commercial oxidation 25 solutions obtained in the commercial oxidation of propylene or isobutylene or by scrubbing the

reaction gases with water.
According to the new process it is also
possible to extract (meth)acrylic acid selectively possible to extract (meth herylic acid selectively 30 from vapors. Thus, the reaction gases obtained in the commercial oxidation of propylene or isobutylene and containing (meth)arrylic acid may be trended with the solvent. Vapors having a high water content are also suitable as

a nga water content are also shades as 35 starting materials. The new process is particu-larly advantageous in the purification of spous containing more than 50% by weight of (meth) actylic acid.

Lactams having 4 to 7 ring members and 40 bearing a hydrocarbon radical with 6 to 20 pearing a hydrocarbon radical with 6 to 20 curbon atoms as substitutent on the airrogen atoms are used as solvents. Suitable compounds, which should be liquid under the reaction conditions, are propiolactums, pyrotidones and oppolations which bear an alkyl, alkenyl, cycloalkyl, aryl or sralkyl radical stayl, alkenyl, cycloalkyl, aryl or sralkyl radical

as substituent on the nitrogen atom. Alkyl radicals with 6 to 20 carbon atoms, cycloalkyl radicals with 6 to 12 carbon atoms, aryl radicals with 6 to 12 carbon atoms and aralkyl radicals

with 7 to 12 carbon atoms, are preferred as hydrocarbon radicals on the nitrogen atom of the lactam. The lactams may also bear alkyl, the factom. The incomes may also bear alkyl, cycloalkyl, anyl or arallyl radicals as substi-tions on one or more, preferably one or two, carbon atoms in the ring. If the lactam bears more than one radical as substituents on carbon atoms, these substituents may be identical or different. Preferred radicals are alkyl groups

having 1 to 6 carbon atoms, cycloalkyl groups having 5 to 8 carbon atoms, aryl groups having 6 to 10 carbon atoms and arallyl groups having 7 to 12 carbon stoms. Examples of sainable compounds are N-ocylpropiolactum, 4-chyl-N-hexylpropiolactum, N-butylpyrroli-

done, 3.5-dimethyl-N-isopropylpyrrollidone, 4, 6-dimethyl-N-enhylpjeridone, N-penylpiperi-done, N-ancylpjeridone, N-burylcaprolatzam and N-octylcaprolatzam. It is particularly advantageous to use pyr-rollidones which bear a hydrocarbon radical

with 6 to 20 carbon atums as substiment on the nitrogen, alkyl and alkenyl groups being preferred as hydrocarbon radicals. Example preferred as hydrocarbon rudicals. Examples are: N-hepstypyrolidane, N-oxylpyrolidane, N-oxylpyrolidane and N-(2-phenylethyl)-pyr-rolidane are also suitable.

In order to avoid difficulties in the distillati purification of the solvents—a measure which is necessary after prolonged enstream periods is accessary are preionged ensurem periods— it is advantageous to use those solvents which contain a total of not more than 25, preferably 7 to 20 and particularly 10 to 17 carbon stoms. to a not particularly to a 11 current since for extracting aqueous (meth)sarylic solutions those solvents are particularly suitable which at room temperature have extremely low water-solubility, i.e. up to about 0.2% by weight, so that the extractant contained in the aqueous solutions can be easily recovered, e.g.

by extraction with an aromatic bydrocarb Obviously mixtures of solvents may be used instead of pure solvents. Mixtures of the abovementioned solvents may be used. It is abovementioned solvenus may be used. It is however also possible to use mixtures of the abovementioned solvenus with other solvenus, such as aliphatic, eyeloaliphatic or aromatic hydrocarbons, halohydrocarbons, ethers or exters; the lactam content should be at least 50% by weight. Many of such mixtures have

lower water solubility than the pure solvents.

The extraction may be carried out within a wide temperature range, between 0° and 200°C particularly between room temperature and 150°C. In the extraction of solutions, however the use of a temperature other than room temperature does not offer any further advan-tages. Extraction is usually carried out at amospheric presure, but superatmospheric pressure, e.g. up to 5 amospheres grage, may also be used. The use of subatmospheric pressure, e.g. 10 to 600 mm Hg, may be

pressure, e.g. 10 to comming, may be advantageous in the extraction of vapors.

The extraction of (meth)scrylic said from solutions is carried out in conventional manner either batchwise or preferably continuously. It is usually carried out in more than one stage. It is advantageous to use continuous extractive columns having 2 or more, particularly 2 to 20 theoretical trays. In many cases, about 95% of the (meth)acrylic acid contained in an

aqueous solution can be separated in 2 stages.
The amount of solvent used is usually 0-1 to 10, advantageously 0.2 to 5, times the amount of the solution, the initial concentration and the desired final concentration being the governing factors. In principle, it is possible to produce

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extracts whose (meth)acrylic acid concentration is as high as the concentration of the squeous solution multiplied by the distribution coefficient. In general, however, solutions of slightly lower concentration are obtained.

5 slightly lower concountration are obtained.

The extraction of (meth)acytic and from vaporens mixtures is also carried out in conventional manner, e.g. by contacting the vaporens with the solvent constructive that the vaporens with the solvent constructive that the vaporens with the solvent constructive that the vaporens with the solvent is supplied to the top of the column. The pressure and the top the column the pressure and the top man the top man to the top of the column. The pressure when the top was said for the other impurities can be written to the contains very little or no (meth)acytic acid. For example, pressures between 100 and 1,000 mm Hg at temperatures of from 50 to 210°C are statuths, for we pressures being used 100°C are statuths, for we pressure being used to 10°C are statuths, for we pressure being used to 10°C are statuths, for we pressure being used to 10°C are statuths, for we pressure being used to 10°C are statuths, for we pressure being used to 10°C are statuths, for which we will be the statuth of the column and upplied to a second to the statuth of the column and upplied to a second to th

bottom of the column and supplied to a second defitilization column where (mechaper)ic acid is 2 distilled off and the solvent is recovered as bottoms product. The olvent is then nevertain sit then expedient in the column of t

regame sometice of accuming it is acrossmental property of the control of the computer and the control of the control of the control column. The mixture of water, (meth)acrylic acid, possibly acric acid and propionic spid, and small amounts of extractant, which mixture is obtained in waper form at the top of the distillation column, is condensed by cooling and supplied to the liquid-liquid extraction column

Concentration of any acetic acid and propionic acid that may be present takes place in the distillation column above the (methlecrylic acid owing to their lower distribution coefficience, i.e. their higher volatility under the

conditions of the centerive distillations, It is a herefore possible to windstayed this layer sensitive states are the form the vapor chamber of many points of the distillation, cohemist where the point of the distillation cohemist where the concentration in the par phase is at a maximum. This sides terms always contains with finalling concentration in the part phase is at a maximum that side of the concentration in the part phase is at a maximum that side of the concentration of the phase is at a maximum value of the phase is at a maximum value of the phase is a concentration of the phase is a side of the phase

which a small mount of unconcer solvent is fed. The wash liquid obtained is improducedly into the distillation column at a suitable point of The (meth)arrylic acid is necovered in a second side stream below the point of which the two samuted carboxylic acids are with

All product streams withdrawn from the 170 vapor chamber contain a small amount of several vapor. In order to two them from solvent they are washed with a reflire of the inquision product, the wash friend being inquision product, the wash friend being recycled to a point in the column at the same, 75 experience of the column at the same of the point at which the stream concerned is with

arawn. In the course of the separation the solvent is cariford with higher-boiling compounds figure crossed as a cream level; is bleed screening a certain twel; is bleed screening solvent is withdrawn, freed from the solvent is withdrawn from the solvent is withdrawn from the solvent is owned by the solvent as owned product and recognition for

In the extraction of a 10 the year of the current o

Salvent	A g of water per 100 g of organic phase	B g of solvent per 100 g of aqueous phass	R
N-isoocrylpyrrolidene N-dedecylpyrrolidene	22 19 20	1·2 0·8 0·9	7-8 4-8 3-14

A denotes the solubility of water in the crysic acid. K is the distribution coefficient organic phase, while B indicates the solubility for scrylic acid.

In the extraction of aqueous solutions contining 10 wr. % of methacrylic acid or 10 wr. % of propionic acid of isobutypic acid or 10 wr. % of propionic acid

		Salvent	Table 4 (Distribution coefficients) K (methacylic acid) (trobutyric acid		K) (propionic acid)
15	٠.	N-isooctylpyrrolidene N-dedecylpyrrolidene	28-5	17-4 10-0	5-6 3-15

Crude aqueous acrylic acid containing 15-14% by weight of acrylic acid, 2-74% by weight of acrylic acid, 2-74% by weight of carbonyl compounds (calculated as acrolich) is used as starting material. N-2-ethylhenyl-pyrrolidone (NOP) is employed as extraction. 1-5 l per hour of aqueous acrylic acid and 15 I per hour of NOP are supplied to the top

and bottom respectively of a pulsed extractive column 40 mm in diameter and having 60 perforated trays, each with 19 holes 2 mm in diameter, the tray spacing being 50 mm. The solution of NOP and acrylic acid which is with-

solution of NOP and actylic acid which is with-forware at the top is peased into a debydration column (1,000 mm in height, 70 mm in diameter, packed with Ruschig rings); a frac-tion consisting mainly of water is obtained at a pressure of 50 mm Hg, a bottoms insuperture 20 of 115°C and a temperature of 35°C at the top and returned to the entractive column. The and returned to the extractive column. The bestumes product obtained in the dehydration column is fed to a point approximately in the middle of a first acotic scale column (2500 nm in height, 40 nm in diameter, packed with straillest-steed wire mesh rings), while at the same time 0'11 per hour of NOI is supplied to the tip of the column. At a pressure of 50 nm

to the tup of the column. At a pressure of 50 mm. Hg, a bottoms temperature of 140°C, a temperature of 47°C at the tup and a reflux ratio of 5: 1 a distillate comaining 186°, by weight of arrylic acid, 70°6, by weight of 5 acctic acid, 108°6, by weight of 6 acctic acid, 108°6, by weight of carbon and 0.75°6, by wright of carbon 10 morpounds is obtained. This distillate is fed to a polant approximatily in the middle of a scond series acid column (J 200 mm. in beight 4.0 mm. in 4 merce.

in the mindle of a second greet acts column (1,200 mm in height, 40 mm in diameter, packed with stainless stud wire mesh rings), while at the same time 0-0151 per hour of NOP is supplied to the top of the column. At w NOP is supplied to the top of the column. As a pressure of 50 mm Hg, a bottoms temperature of 70°C, a temperature of 46°C at the top and a reflux ratio of 5°. I a distillate is obtained which contains 80% by weight of acrtic acid, 15-6% by weight of warr and 0.55% by weight. of carbonyl compounds. The amount of sectic acid contained in this distillate is 83-3%

accus cana contained in this distillate is 83-3%, of the amount originally contained in the squoous solution. The bottoms product from this column is recycled to the first accele and column. It contains 72%, by weight of explicit acid, 10%, by weight of accele acid, 10% by weight of accele acid, 10% of the column of the co

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The boitems product from the first accided to the upper part of an acrylic acid column (1,200 mm in height, 40 mm in diameter, packed with stainless steel wire mesh rings). At a reflux ratio of 1: 1, a 60 pressure of 30 mm Hg, a bottoms temperature of 170°C and a temperature of 66°C at the top acrylic acid having a purity of about 976 to 98% by weight (96% of the amount originally contained in the aqueous solution) is obtained

The aqueous phase obtained in the extrac tion column contains 0.4% by weight of acrylic acid and acetic acid and 2.98% by

weight of carbonyl compounds.

The NOP obtained as bottoms product in the acrylic acid column is recycled, a bleed stream of about 25% being previously purified

distillation. WHAT WE CLAIM IS:-

 In a process for separating methacrylic scid and/or acrylic acid from mixtures conacin annors activitie acid more acrylic acid the imigm methacrylic acid andors acrylic acid the improvement which comprises extracting said mixtures with a lactam having 4 to 7 ring members and bearing a hydrocarbon radical with 6 to 20 carbon atoms as substituent on the nitrogen atom as extractant.

2. A process as claimed in claim I wherein a prolidone bearing an alkyl radical with 6 to 20 carbon atoms or an alkenyl radical with 6 to 20 carbon atoms as substituent on the nitrogen

20 carbon atoms is substituted to the nurogen atoms is used as extractant.

3. A process as claimed in claim 2 for separating methacrytic acid and/or acrytic acid 90 from equeous solutions containing 2 to 50%, by weight of methacrytic acid and/or acrytic acid comprising extracting said solutions at a temperature of from 0° to 200°C and a pressure om 10 mm Hg to 5 atmospheres gas

distilling off the methacrylic acid and/or symbolytic scrid and/or acrylic acid in said perfectly and recycling the pyrrolidone and recycling the pyrrolidone bained as lottoms product in the distillation 100

4. A process for separating scrylic acid from a mixture containing scrylic acid substantially as described in the foregoing Example.

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